

STIC Search Report Biotech-Chem Library

STIC Database Tracking Number

TO: Ruth Davis

Location: REM-3D71

Art Unit: 1651

Thursday, February 12, 2004

Case Serial Number: 09/284806

From: Alex Waclawiw

Location: Biotech-Chem Library

Rem 1A71

Phone: 308-4491

Alexandra.waclawiw@uspto.gov

Search Notes

Examiner Davis,

This is the structure for the compound that you specified. The inventor named it differently than CAS. Ignore the stereochemistry since a basic structure search includes all stereochemistry. See references for the structure hits.

If you have any questions please feel free to contact me.

Alexandra Waclawiw

Me Me Me Me Me Me
$$(CH_2)_5$$
 S E E E O



clenuic and recimical information center

Patent Intranet > NPL Virtual Library > Request a Prior Art Search

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114157

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Tech Center:	
© TC 1600	
Enter your Contact Information below:	RECEIVED FEB 11 200
Name: Ruth Davis	
Employee Number: 78220 Phone: 571-272-0915	VED 200
Art Unit or Office: 1651 Building & Room Number: REM 3D71	
Enter the case serial number (Required): 09/284,806 If not related to a patent application, please enter NA here.	<u>.</u>
Class / Subclass(es)	
Earliest Priority Filing Date: 10/21/97	
Format preferred for results: Paper Diskette E-mail	0-
Provide detailed information on your search topic:	48

- In your own words, describe in detail the concepts or subjects you want us to search.
- Include synonyms, keywords, and acronyms. Define terms that have special meanings.
- *For Chemical Structure Searches Only*
 Include the elected species or structures, keywords, synonyms, acronyms, and registry
 numbers
- *For Sequence Searches Only*
 Include all pertinent information (parent, child, divisional, or issued patent numbers) along with the appropriate serial number.
- *For Foreign Patent Family Searches Only* Include the country name and patent number.
- Provide examples or give us relevant citations, authors, etc., if known.
- FAX or send the abstract, pertinent claims (not all of the claims), drawings, or chemical structures to your EIC or branch library.

25. 50

Enter your Search Topic Information below:

Claims I and 3.

Specifically:

3-(C55,65)-5,6-dihydro-5-(c65)-4,6-dimethydodeca2E,4E-diencyl)-2H-pyran-2-on:-6-yl)
prop-2E-enoic acid

Special Instructions and Other Comments:

(For fastest service, let us know the best times to contact you, in case the searcher needs further clarification on your search.)

Press ALT + F, then P to print this screen for your own information.

SEND RESET

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Last Modified: 12/05/2003 15:08:46

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=> d his
```

(FILE 'REGISTRY' ENTERED AT 12:15:04 ON 12 FEB 2004)
DEL HIS Y

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FILE 'REGISTRY' ENTERED AT 12:16:43 ON 12 FEB 2004
ACT DAVISCLM1/A
```

```
STR
L1
            58) SEA FILE=REGISTRY SSS FUL L1
   (
L2
L3
              STR
             6 SEA FILE=REGISTRY SUB=L2 SSS FUL L3
L4
               ACT FORMII/A
              -----
L5
              STR
            58) SEA FILE=REGISTRY SSS FUL L5
L6 (
          4085) SEA FILE=REGISTRY ABB=ON PLU=ON C8H10O3
L7 (
             4) SEA FILE=REGISTRY ABB=ON PLU=ON L7 AND L6
L8 (
           363)SEA FILE=REGISTRY ABB=ON PLU=ON C8H8O5
L9 (
            1) SEA FILE=REGISTRY ABB=ON PLU=ON L9 AND L6
L10 (
             5 SEA FILE=REGISTRY ABB=ON PLU=ON L8 OR L10
L11
              _____
               ACT FORMIIIA/A
              -----
          3569) SEA FILE=REGISTRY ABB=ON PLU=ON C14H24O2
L12 (
            12) SEA FILE=REGISTRY ABB=ON PLU=ON L12 AND 2 4 DODECADIENOIC
L13 (
             3 SEA FILE=REGISTRY ABB=ON PLU=ON L13 AND 4 6 DIMETHYL
L14
              _____
               SELECT RN L14 1-3
               SELECT RN L16 1-3
               SET SMARTSELECT ON
           SEL L11 1- RN : 5 TERMS
L15
               SET SMARTSELECT OFF
             0 S L15/CRN
L16
               SET SMARTSELECT ON
           SEL L14 1- RN : 3 TERMS
L17
               SET SMARTSELECT OFF
             0 S L17/CRN
L18
     FILE 'CAPLUS' ENTERED AT 12:25:57 ON 12 FEB 2004
             6 S L4
L19
             2 S L11 AND L14
L20
```

=> fil reg FILE 'REGISTRY' ENTERED AT 12:28:49 ON 12 FEB 2004 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2004 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 11 FEB 2004 HIGHEST RN 649538-27-2 DICTIONARY FILE UPDATES: 11 FEB 2004 HIGHEST RN 649538-27-2

TSCA INFORMATION NOW CURRENT THROUGH JULY 14, 2003

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

VAR G1=CO2H/ME NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:
RSPEC I
NUMBER OF NODES IS

STEREO ATTRIBUTES: NONE

L2 (58)SEA FILE=REGISTRY SSS FUL L1
L3 STR

Page 2 searched by Alex Waclawiw

VAR G1=CO2H/ME NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 26

STEREO ATTRIBUTES: NONE

6 SEA FILE=REGISTRY SUB=L2 SSS FUL L3

100.0% PROCESSED

6 ITERATIONS

6 ANSWERS

SEARCH TIME: 00.00.01

=> d que 111

STR

VAR G1=CO2H/ME NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 11

STEREO ATTRIBUTES: NONE

58) SEA FILE=REGISTRY SSS FUL L5 L6 (4085) SEA FILE=REGISTRY ABB=ON PLU=ON C8H10O3 L74) SEA FILE=REGISTRY ABB=ON PLU=ON L7 AND L6 L8 363) SEA FILE=REGISTRY ABB=ON PLU=ON C8H8O5 L9 1) SEA FILE=REGISTRY ABB=ON PLU=ON L9 AND L6 L10 (5 SEA FILE=REGISTRY ABB=ON PLU=ON L8 OR L10 L11

=> d que stat l14

L12 (3569) SEA FILE=REGISTRY ABB=ON PLU=ON C14H24O2

12) SEA FILE=REGISTRY ABB=ON PLU=ON L12 AND 2 4 DODECADIENOIC L13 (

3 SEA FILE=REGISTRY ABB=ON PLU=ON L13 AND 4 6 DIMETHYL L14 .

=> d his 115-118

(FILE 'REGISTRY' ENTERED AT 12:16:43 ON 12 FEB 2004)

SELECT RN L14 1-3 SELECT RN L16 1-3

SET SMARTSELECT ON

SEL L11 1- RN : 5 TERMS L15

Page 3 searched by Alex Waclawiw

Davis 09/284,806

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SET SMARTSELECT OFF

L16 0 S L15/CRN
SET SMARTSELECT ON

L17 SEL L14 1- RN : 3 TERMS
SET SMARTSELECT OFF

L18 0 S L17/CRN
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=> fil caplus FILE 'CAPLUS' ENTERED AT 12:30:01 ON 12 FEB 2004 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 12 Feb 2004 VOL 140 ISS 7 FILE LAST UPDATED: 11 Feb 2004 (20040211/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

'OBI' IS DEFAULT SEARCH FIELD FOR 'CAPLUS' FILE

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=> d que nos 119
1.1
                STR
          58) SEA FILE=REGISTRY SSS FUL L1
L_2
L3
                STR
             6 SEA FILE=REGISTRY SUB=L2 SSS FUL L3
L4
              6 SEA FILE=CAPLUS ABB=ON PLU=ON L4
L19
=> d que nos 120
L5
                STR
             58) SEA FILE=REGISTRY SSS FUL L5
L6
          4085) SEA FILE=REGISTRY ABB=ON PLU=ON C8H10O3
L7
            4) SEA FILE=REGISTRY ABB=ON PLU=ON L7 AND L6
L_8
           363) SEA FILE=REGISTRY ABB=ON PLU=ON C8H8O5
L9
             1) SEA FILE=REGISTRY ABB=ON PLU=ON L9 AND L6
L10 (
             5 SEA FILE=REGISTRY ABB=ON PLU=ON L8 OR L10
L11
           3569) SEA FILE=REGISTRY ABB=ON
                                         PLU=ON C14H24O2
L12 (
                                        PLU=ON L12 AND 2 4 DODECADIENOIC
            12) SEA FILE=REGISTRY ABB=ON
L13 (
             3 SEA FILE=REGISTRY ABB=ON PLU=ON L13 AND 4 6 DIMETHYL
L14
              2 SEA FILE=CAPLUS ABB=ON PLU=ON L11 AND L14
L20
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=> d .ca hitstr 119 1-6;d .ca hitstr 120 1-2

L19 ANSWER 1 OF 6 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2002:340575 CAPLUS

DOCUMENT NUMBER: 137:108320

TITLE: Identifying protein kinase inhibitors using an assay

Page 4 searched by Alex Waclawiw

Davis 09/284,806

based on inhibition of aerial hyphae formation in

Streptomyces

AUTHOR(S): Waters, Barbara; Saxena, Geeta; Wanggui, Yangsheng;

Kau, David; Wrigley, Stephen; Stokes, Richard; Davies,

Julian

CORPORATE SOURCE: Cubist Pharmaceuticals, Inc., Vancouver, BC, V6T 1Z3,

Can.

SOURCE: Journal of Antibiotics (2002), 55(4), 407-416

CODEN: JANTAJ; ISSN: 0021-8820

PUBLISHER: Japan Antibiotics Research Association

DOCUMENT TYPE: Journal LANGUAGE: English

We have identified a strain of Streptomyces in which aerial hyphae formation appears to be especially sensitive to inhibition by protein kinase inhibitors. Using this assay, a number of bacterial cultures have been screened and novel inhibitors of eukaryotic protein kinases have been identified. Since M. tuberculosis possesses multiple eukaryotic-like protein kinase genes, we tested the active kinase inhibitors for the inhibition of mycobacterial growth and obtained several potent compds. This identifies a new biochem. class of antimycobacterial agents.

CC 16-1 (Fermentation and Bioindustrial Chemistry)

Section cross-reference(s): 9

24730-31-2, Surfactin 18791-21-4, Pyridomycin 446-72-0, Genistein 71897-07-9, Ag-1295 131956-33-7, Depsidomycin 27127-62-4, Viscosin 169062-92-4, Cyclomarin a 133550-30-8, Ag-490 153436-53-4, Ag-1478 192819-12-8, XR 587 **207220-91-5**, Xr-379 339320-58-0, Xr-774 405149-80-6, Xr-543 405149-83-9, Xr-318 343780-48-3, Xr-336 405150-10-9, Xr-315 405150-11-0, Xr-475 405150-05-2, Xr-665 405150-13-2, Xr-819

RL: BSU (Biological study, unclassified); BIOL (Biological study) (identifying protein kinase inhibitors using assay based on inhibition of aerial hyphae formation in Streptomyces)

IT 207220-91-5, Xr-379

RL: BSU (Biological study, unclassified); BIOL (Biological study) (identifying protein kinase inhibitors using assay based on inhibition of aerial hyphae formation in Streptomyces)

RN 207220-91-5 CAPLUS

CN 2,4-Dodecadienoic acid, 4,6-dimethyl-, (2S,3S)-2-[(1E)-2-carboxyethenyl]-3,6-dihydro-6-oxo-2H-pyran-3-yl ester, (2E,4E,6S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry as shown.

Me Me
$$(CH_2)_5$$
 S E CO_2H

REFERENCE COUNT:

40 THERE ARE 40 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

Davis 09/284,806

```
ANSWER 2 OF 6 CAPLUS COPYRIGHT 2004 ACS on STN
                        2002:220383 CAPLUS
ACCESSION NUMBER:
                         136:259921
DOCUMENT NUMBER:
                         Antibacterial agents and methods of identification
TITLE:
                         Davies, Julian E.; Waters, Barbara
INVENTOR(S):
                         Cubist Pharmaceuticals, Inc., USA
PATENT ASSIGNEE(S):
                         PCT Int. Appl., 41 pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
                         English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                           APPLICATION NO. DATE
                      KIND DATE
     PATENT NO.
     _____
                     ----
                            20020321
                                           WO 2001-US28913 20010917
     WO 2002022138
                      Α1
     WO 2002022138
                      C2
                           20031106
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
            LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PH, PL,
             PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG,
             US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     AU 2001096253
                      A5 20020326
                                          AU 2001-96253
                                                            20010917
                                        US 2000-233004P P 20000915
PRIORITY APPLN. INFO.:
                                        WO 2001-US28913 W 20010917
     The present invention provides an assay based on Streptomyces species in
     which aerial hyphae formation and sporulation appear to be especially sensitive
     to inhibition by protein kinase inhibitors which are also antibacterial
     agents. Using this Streptomyces-based assay and a growth inhibitory
     assay, a number of bacterial cultures have been examined and several potential
     novel inhibitors of antimycobacterial agents have been identified. The
     antibacterial screening method of the invention comprises two step: (a)
     contacting a growing culture of Streptomyces griseus or Streptomyces 85E
     with a test compound for a time sufficient to allow the test compound to alter
     aerial mycelial development or sporulation, and (b) contacting
     mycobacterium cells with the test compound of step (a) for a time sufficient
     to allow the test compound to inhibit growth of the mycobacterium. Test
     compds. that tested pos. in both step (a) and step (b) are antibacterial
     compds. of the invention.
     ICM A61K031-70
TC
     ICS A61K031-33; A61K035-00; C12Q001-48
     10-5 (Microbial, Algal, and Fungal Biochemistry)
CC
     Section cross-reference(s): 1
     207220-91-5, XR 379
TT
     RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (XR 379; antibacterial agents and methods of identification based on
        Streptomyces species and inhibition of Mycobacterium and purification from
        microbial cell culture supernatant)
     207220-91-5, XR 379
IT
     RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (XR 379; antibacterial agents and methods of identification based on
        Streptomyces species and inhibition of Mycobacterium and purification from
        microbial cell culture supernatant)
RN
     207220-91-5 CAPLUS
```

2,4-Dodecadienoic acid, 4,6-dimethyl-, (2S,3S)-2-[(1E)-2-carboxyethenyl]-3,6-dihydro-6-oxo-2H-pyran-3-yl ester, (2E,4E,6S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

Me Me
$$CCH_2$$
 S S E CO_2H

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 1 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L19 ANSWER 3 OF 6 CAPLUS COPYRIGHT 2004 ACS on STN 2001:224151 CAPLUS

ACCESSION NUMBER:

135:32775

DOCUMENT NUMBER: TITLE:

Scale-up of filamentous organisms from tubes and

shake-flasks into stirred vessels

AUTHOR (S):

Katzer, Werner; Blackburn, Mark; Charman, Kevin;

Martin, Steven; Penn, Julia; Wrigley, Stephen

CORPORATE SOURCE:

TerraGen Discovery (UK) Ltd., Slough, UK

SOURCE:

Biochemical Engineering Journal (2001), 7(2), 127-134

CODEN: BEJOFV; ISSN: 1369-703X

PUBLISHER:

Elsevier Science S.A.

DOCUMENT TYPE:

Journal LANGUAGE: English

The choice of small-scale fermentation systems contributes significantly to a successful scale-up. Creasing of flasks and the chosen shaker parameters influence the production of secondary metabolites in a strain- and even compound-specific manner. Using actinomycetes and fungi as model organisms the influence of the small-scale fermentation system on the production of various

secondary metabolites is described and the effects on screening success and scale-up are considered.

16-2 (Fermentation and Bioindustrial Chemistry) CC

IT 74720-35-7P, xr334 102228-99-9P, diastovaricin II 157842-16-5P, XR 368 207225-51-2P, XR 573 343780-48-3P 207220-91-5P, XR 379

RL: BMF (Bioindustrial manufacture); BPN (Biosynthetic preparation); BIOL (Biological study); PREP (Preparation)

(scale-up of filamentous organisms from tubes and shake-flasks into stirred vessels)

207220-91-5P, XR 379 ΙT

RL: BMF (Bioindustrial manufacture); BPN (Biosynthetic preparation); BIOL (Biological study); PREP (Preparation)

(scale-up of filamentous organisms from tubes and shake-flasks into stirred vessels)

207220-91-5 CAPLUS RN

2,4-Dodecadienoic acid, 4,6-dimethyl-, (2S,3S)-2-[(1E)-2-carboxyethenyl]-CN3,6-dihydro-6-oxo-2H-pyran-3-yl ester, (2E,4E,6S)- (9CI) (CA INDEX NAME) Absolute stereochemistry.
Double bond geometry as shown.

REFERENCE COUNT:

THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L19 ANSWER 4 OF 6 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

1999:720265 CAPLUS

DOCUMENT NUMBER:

131:310551

TITLE:

Preparation of dihydropyrones as cytokine production

inhibitors.

INVENTOR(S):

Hayes, Martin Alistair; Hardick, David James; Tang, Jenny Seukgin; Ryder, Hamish; Folkes, Adrian John;

Tatsuoka, Toshio; Matsui, Masashi Xenova Limited, UK; Suntory Limited Brit. UK Pat. Appl., 59 pp., 59 pp.

CODEN: BAXXDU

SOURCE:

Patent

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT ASSIGNEE(S):

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE		
GB 2336362	A1	19991020	GB 1998-8196	19980417		
JP 11335365	A2	19991207	JP 1999-109982	19990416		
US 6197811	B1	20010306	US 1999-292961	19990416		
PRIORITY APPLN. INFO.	:	GB	1998-8196 A	19980417		
OTHER SOURCE(S):	MΑ	RPAT 131:310551				
CT	,					

Title compds. [I; X = O, NH; R1 = R3CO, ArCH2, R5OCH2; R3 = R4CH:CH, alkyl, (alkyl-substituted) fluorenyl, oxofluorenyl; R4 = alkyl, alkenyl, aryl, unsatd. heterocyclyl; Ar = aryl; R5 = alkyl optionally interrupted by 1-2 O; R2 = Me, R6O2C; R6 = alkyl], were prepared Thus, di-Et azodicarboxylate in THF was added dropwise to a suspension of phomalactone, Ph3P, and PhCO2H followed by stirring overnight to give 43%

[(2S,3R)-3,6-dihydro-6-oxo-(2E-prop-1-enyl)-2H-pyran-3-yl]benzoate. The latter inhibited IL-1 β release from human peripheral blood mononuclear cells with IC50 = 2.4 μ M.

- IC ICM C07D309-30 ICS A61K031-365
- CC 27-13 (Heterocyclic Compounds (One Hetero Atom))

Section cross-reference(s): 1

TT 247188-29-0P 247188-30-3P 247188-31-4P 247188-32-5P 247188-33-6P 247188-35-8P 247188-36-9P 247188-37-0P 247188-38-1P 247188-34-7P 247188-41-6P 247188-42-7P 247188-39-2P **247188-40-5P** 247188-46-1P 247188-47-2P 247188-44-9P 247188-45-0P 247188-43-8P 247188-48-3P 247188-49-4P 247188-50-7P 247188-51-8P 247188-52-9P 247188-55-2P 247188-56-3P 247188-57-4P 247188-53-0P 247188-54-1P 247188-60-9P 247188-61-0P 247188-58-5P 247188-59-6P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of dihydropyrones as cytokine production inhibitors) 65-85-0, Benzoic acid, reactions 71-23-8, Propanol, reactions 1-Butanol, reactions 92-92-2, 4-Biphenylcarboxylic acid 98-88-4, 100-39-0, Benzyl bromide 100-52-7, Benzaldehyde, Benzoyl chloride 784-50-9, 9-Fluorenone-2-carboxylic acid 1573-92-8, reactions 9-Fluorenone-1-carboxylic acid 3218-36-8, 4-Biphenylcarboxaldehyde 3970-21-6, 2-Methoxyethoxymethyl chloride 5728-52-9, 4-Biphenylacetic 6276-03-5, 1-Fluorenecarboxylic acid 7071-83-2, 5731-13-5 14002-51-8, 4-Biphenylcarbonyl chloride 9-Fluorenone-4-carbonyl chloride 15690-24-1 28921-94-0, Phomalactone 30084-90-3, 2-32466-54-9, trans-2-Dodecenoic acid 73373-17-8, Fluorenecarboxaldehyde 138875-82-8 **207220-91-5** 4-Iodomethylbiphenyl RL: RCT (Reactant); RACT (Reactant or reagent)

(preparation of dihydropyrones as cytokine production inhibitors)

247188-40-5P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of dihydropyrones as cytokine production inhibitors)

RN 247188-40-5 CAPLUS

TT

CN 2,4-Dodecadienoic acid, 4,6-dimethyl-, (2S,3R)-3,6-dihydro-6-oxo-2-(1E)-1-propenyl-2H-pyran-3-yl ester, (2E,4E,6S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

IT 207220-91-5

RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation of dihydropyrones as cytokine production inhibitors)

207220-91-5 CAPLUS RN

2,4-Dodecadienoic acid, 4,6-dimethyl-, (2S,3S)-2-[(1E)-2-carboxyethenyl]-CN3,6-dihydro-6-oxo-2H-pyran-3-yl ester, (2E,4E,6S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

L19 ANSWER 5 OF 6 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

1999:709567 CAPLUS

DOCUMENT NUMBER:

132:61362

TITLE:

A novel (6S)-4,6-dimethyldodeca-2E,4E-dienoyl ester of

phomalactone and related α -pyrone esters from a Phomopsis sp. with cytokine production inhibitory

activity

AUTHOR (S):

Wrigley, Stephen K.; Sadeghi, Roya; Bahl, Sangeeta; Whiting, Andrew J.; Ainsworth, A. Martyn; Martin, Steven M.; Katzer, Werner; Ford, Robert; Kau, David A.; Robinson, Neil; Hayes, Martin A.; Elcock, Claire;

Mander, Thomas; Moore, Michael

CORPORATE SOURCE:

SOURCE:

TerraGen Discovery (UK) Ltd., Slough, SL1 4EQ, UK Journal of Antibiotics (1999), 52(10), 862-872

CODEN: JANTAJ; ISSN: 0021-8820

Japan Antibiotics Research Association

PUBLISHER:

DOCUMENT TYPE:

Journal English

LANGUAGE:

GΙ

Me Me
H

$$R = Me$$
 $R = Me$
 $R = Me$
 $R = CO_2H$

Me Me
$$_{\rm OH}$$
 $_{\rm OH}$ $_{\rm HO_2C}$ $_{\rm H}$ $_{\rm OO}$ $_{\rm III}$

A series of novel 6-substituted 5,6-dihydro-5-hydroxy- α -pyrone AB esters isolated from fermns. of a Phomopsis sp. (Xenova culture collection number X22502) have been identified as inhibitors of lipopolysaccharide (LPS) - induced cytokine production These include the phomalactone (6S)-4,6-dimethyldodecadien-2E,4E-dienoyl ester (I), and two analogs (II and III) bearing a prop-2E-enoic acid moiety at the 6-position of the α -pyrone ring. (6S)-4,6-Dimethyl-2E,4E-dienoic acid and a hydroxylated analog were also isolated and characterized. The most potent cytokine production inhibitor was I, which inhibited LPS-induced tumor necrosis factor α (TNF α) production by U937 cells and LPS-induced interleukin 1β (IL- 1β) production by peripheral blood mononuclear cells (PBMC) with IC50 values of 80 nM and 190 nM resp. The effect of I in PBMC was selective for IL-1 β relative to TNF α . The inhibition of IL-1 β production by I involved a post-translational mechanism of action at the level of $IL-1\beta$ secretion as demonstrated by the lack of an effect on cell-associated IL-1 β production I showed no effect on the activity of caspase 1 in cytosolic exts. from the THP1 monocytic cell line.

CC 10-1 (Microbial, Algal, and Fungal Biochemistry)
Section cross-reference(s): 1

IT 138875-82-8P 207220-91-5P 207220-92-6P 253351-45-0P 253351-46-1P

RL: BAC (Biological activity or effector, except adverse); BOC (Biological occurrence); BPN (Biosynthetic preparation); BSU (Biological study, unclassified); PUR (Purification or recovery); THU (Therapeutic use); BIOL (Biological study); OCCU (Occurrence); PREP (Preparation); USES (Uses) (novel phomalactone dimethyldodecadienoyl ester and related pyrone esters from Phomopsis with cytokine production inhibitory activity)

207220-91-5P 207220-92-6P 253351-45-0P
RL: BAC (Biological activity or effector, except adverse); BOC (Biological occurrence); BPN (Biosynthetic preparation); BSU (Biological study, unclassified); PUR (Purification or recovery); THU (Therapeutic use); BIOL (Biological study); OCCU (Occurrence); PREP (Preparation); USES (Uses) (novel phomalactone dimethyldodecadienoyl ester and related pyrone esters from Phomopsis with cytokine production inhibitory activity)

TT

RN 207220-91-5 CAPLUS

CN 2,4-Dodecadienoic acid, 4,6-dimethyl-, (2S,3S)-2-[(1E)-2-carboxyethenyl]-3,6-dihydro-6-oxo-2H-pyran-3-yl ester, (2E,4E,6S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

RN 207220-92-6 CAPLUS

CN 2,4-Dodecadienoic acid, 4,6-dimethyl-, (2S,3S)-3,6-dihydro-6-oxo-2-(1E)-1-propenyl-2H-pyran-3-yl ester, (2E,4E,6S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry as shown.

Me Me
$$(CH_2)_5$$
 S E E O O

RN 253351-45-0 CAPLUS

CN 2,4-Dodecadienoic acid, 11-hydroxy-4,6-dimethyl-, (2S,3S)-2-[(1E)-2-carboxyethenyl]-3,6-dihydro-6-oxo-2H-pyran-3-yl ester, (2E,4E)- (9CI) (CFINDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

OH Me Me
$$E = E$$
 CO_2H CO_2H

REFERENCE COUNT:

THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS 22 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

CAPLUS COPYRIGHT 2004 ACS on STN L19 ANSWER 6 OF 6

ACCESSION NUMBER: DOCUMENT NUMBER:

1998:268500 CAPLUS

128:317258

TITLE:

5,6-Dihydro- α -pyrone cytokine production

inhibitors, their production and preparation, and

their therapeutic use

INVENTOR(S):

Wrigley, Stephen Keith; Bahl, Sangeeta; Guilani, Roya Mansour Sadeghi; Moore, Michael; Katzer, Werner Albert; Martin, Steven Michael; Kau, David Andrew;

Whiting, Andrew Jonathan; Robinson, Neil; Hayes,

Martin Alistair; Mander, Thomas Haydn

PATENT ASSIGNEE(S):

Xenova Ltd., UK; Wrigley, Stephen Keith; Bahl,

Sangeeta; Guilani, Roya Mansour Sadeghi; Moore, Michael; Katzer, Werner Albert; Martin, Steven Michael

PCT Int. Appl., 37 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

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WO	9817	661		Α.	1	1998	0430		W(199	97-GI	3290	7	1997	1021		
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														MW,			
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	UA,	UG,
														ТJ,			
	RW:	GH,	KE,	LS,	MW,	SD,	SZ,	UG,	ZW,	ΑT,	BE,	CH,	DE,	DK,	ES,	FΙ,	FR,
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		GN,	ML,	MR,	NE,	SN,	TD,	TG									
AU	9747	154		A1 19980515				AU 1997-47154				19971021					
GB	2333	294		A	1	1999	0721		G1	B 19	99-8	524		1997	1021		
GB	2333	294		B:	2 .,	2000	1018										
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								WO 1997-GB2907 W					W	1997	1021		

GΙ

- AB A 5,6-dihydro- α -pyrone I (R = CO2H or CH3 and R1, R2 = H; or R = CO2H and one of R1 and R2 is H and the other is OH; or when R is CO2H, a pharmaceutically or veterinarily acceptable salt thereof) are provided. Processes for producing these compds., and their use as cytokine production inhibitors, are also described. Fermentative production and preparative esterification are included.
- IC ICM C07D309-32 ICS C12P007-40; C12P017-06; C07C059-42; C07C057-03; C12N001-14; C12N001-14; C12R001-645; C12R001-79
- CC 1-7 (Pharmacology)
 Section cross-reference(s): 16, 27, 63
- 207220-92-6P
 RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PUR (Purification or recovery); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 - $(5,6\text{-dihydro-}\alpha\text{-pyrone cytokine production inhibitors, production and preparation, and therapeutic use)$
- 207220-91-5P 207220-93-7P 207220-94-8P

 RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PUR (Purification or recovery); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 - $(5,6\text{-dihydro-}\alpha\text{-pyrone}$ cytokine production inhibitors, production and preparation, and therapeutic use)
 - 207220-92-6P
 RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PUR (Purification or recovery); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 - $(5,6\text{-dihydro-}\alpha\text{-pyrone cytokine production inhibitors, production and preparation, and therapeutic use)$
- RN 207220-92-6 CAPLUS CN 2,4-Dodecadienoic acid, 4,6-dimethyl-, (2S,3S)-3,6-dihydro-6-oxo-2-(1E)-1-propenyl-2H-pyran-3-yl ester, (2E,4E,6S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

TT

IT 207220-91-5P 207220-93-7P 207220-94-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PUR (Purification or recovery); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (5,6-dihydro- α -pyrone cytokine production inhibitors, production and preparation, and therapeutic use)

RN 207220-91-5 CAPLUS

CN 2,4-Dodecadienoic acid, 4,6-dimethyl-, (2S,3S)-2-[(1E)-2-carboxyethenyl]-3,6-dihydro-6-oxo-2H-pyran-3-yl ester, (2E,4E,6S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

RN 207220-93-7 CAPLUS

CN 2,4-Dodecadienoic acid, 9-hydroxy-4,6-dimethyl-, (2S,3S)-2-[(1E)-2-carboxyethenyl]-3,6-dihydro-6-oxo-2H-pyran-3-yl ester, (2E,4E,6S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

207220-94-8 CAPLUS RN

2,4-Dodecadienoic acid, 11-hydroxy-4,6-dimethyl-, (2S,3S)-2-[(1E)-2-ÇN carboxyethenyl]-3,6-dihydro-6-oxo-2H-pyran-3-yl ester, (2E,4E,6S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

CAPLUS COPYRIGHT 2004 ACS on STN L20 ANSWER 1 OF 2

2

ACCESSION NUMBER:

1999:720265 CAPLUS

DOCUMENT NUMBER:

131:310551

TITLE:

Preparation of dihydropyrones as cytokine production

inhibitors.

INVENTOR (S):

Hayes, Martin Alistair; Hardick, David James; Tang,

Jenny Seukgin; Ryder, Hamish; Folkes, Adrian John;

Tatsuoka, Toshio; Matsui, Masashi

PATENT ASSIGNEE(S): SOURCE:

Xenova Limited, UK; Suntory Limited Brit. UK Pat. Appl., 59 pp., 59 pp.

CODEN: BAXXDU

DOCUMENT TYPE:

Patent

LANGUAGE:...

FAMILY ACC. NUM. COUNT:

English

PATENT INFORMATION:

PATENT NO.

KIND DATE APPLICATION NO. DATE

Page 16 searched by Alex Waclawiw

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GB 1998-8196
     GB 2336362
                        A1
                              19991020
                                                                19980417
                                              JP 1999-109982
     JP 11335365
                        A2
                              19991207
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                                              US 1999-292961
     US 6197811
                        B1
                              20010306
                                                                19990416
                                          GB 1998-8196
                                                               19980417
PRIORITY APPLN. INFO .:
                          MARPAT 131:310551
OTHER SOURCE(S):
GI
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$$\mathbb{R}^{1\chi}$$
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Title compds. [I; X = O, NH; R1 = R3CO, ArCH2, R5OCH2; R3 = R4CH:CH, alkyl, (alkyl-substituted) fluorenyl, oxofluorenyl; R4 = alkyl, alkenyl, aryl, unsatd. heterocyclyl; Ar = aryl; R5 = alkyl optionally interrupted by 1-2 O; R2 = Me, R6O2C; R6 = alkyl], were prepared Thus, di-Et azodicarboxylate in THF was added dropwise to a suspension of phomalactone, Ph3P, and PhCO2H followed by stirring overnight to give 43% [(2S,3R)-3,6-dihydro-6-oxo-(2E-prop-1-enyl)-2H-pyran-3-yl]benzoate. The latter inhibited IL-1 β release from human peripheral blood mononuclear cells with IC50 = 2.4 μ M.

IC ICM C07D309-30 ICS A61K031-365

CC 27-13 (Heterocyclic Compounds (One Hetero Atom))

Section cross-reference(s): 1

71-23-8, Propanol, reactions 65-85-0, Benzoic acid, reactions TΤ 1-Butanol, reactions 92-92-2, 4-Biphenylcarboxylic acid 98-88-4. 100-52-7, Benzaldehyde, Benzoyl chloride 100-39-0, Benzyl bromide 784-50-9, 9-Fluorenone-2-carboxylic acid 1573-92-8, reactions 3218-36-8, 4-Biphenylcarboxaldehyde 9-Fluorenone-1-carboxylic acid 3970-21-6, 2-Methoxyethoxymethyl chloride 5728-52-9, 4-Biphenylacetic 7071-83-2, 5731-13-5 - 6276-03-5; 1-Fluorenecarboxylic acid 9-Fluorenone-4-carbonyl chloride 14002-51-8, 4-Biphenylcarbonyl chloride 15690-24-1 **28921-94-0**, Phomalactone 30084-90-3, 2-Fluorenecarboxaldehyde 32466-54-9, trans-2-Dodecenoic acid 73373-17-8, 4-Iodomethylbiphenyl **138875-82-8** 207220-91-5 RL: RCT (Reactant); RACT (Reactant or reagent)

(preparation of dihydropyrones as cytokine production inhibitors)

IT **107741-12-8P** 247188-62-1P 247188-63-2P 247188-64-3P 247188-65-4P 247188-66-5P 247188-67-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of dihydropyrones as cytokine production inhibitors)

IT 28921-94-0, Phomalactone 138875-82-8

RL: RCT (Reactant); RACT (Reactant or reagent)

(preparation of dihydropyrones as cytokine production inhibitors)

RN 28921-94-0 CAPLUS

CN 2H-Pyran-2-one, 5,6-dihydro-5-hydroxy-6-(1E)-1-propenyl-, (5S,6S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

RN 138875-82-8 CAPLUS

CN 2,4-Dodecadienoic acid, 4,6-dimethyl-, (2E,4E,6S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry as shown.

IT 107741-12-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of dihydropyrones as cytokine production inhibitors)

RN 107741-12-8 CAPLUS

CN 2H-Pyran-2-one, 5,6-dihydro-5-hydroxy-6-(1E)-1-propenyl-, (5R,6S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-). Double bond geometry as shown.

L20 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

1998:268500 CAPLUS

DOCUMENT NUMBER: TITLE:

128:317258 5,6-Dihydro- α -pyrone cytokine production

inhibitors, their production and preparation, and

their therapeutic use

INVENTOR(S):

Wrigley, Stephen Keith; Bahl, Sangeeta; Guilani, Roya Mansour Sadeghi; Moore, Michael; Katzer, Werner Albert; Martin, Steven Michael; Kau, David Andrew; Whiting, Andrew Jonathan; Robinson, Neil; Hayes,

Martin Alistair; Mander, Thomas Haydn

PATENT ASSIGNEE(S):

Xenova Ltd., UK; Wrigley, Stephen Keith; Bahl, Sangeeta; Guilani, Roya Mansour Sadeghi; Moore,

Michael; Katzer, Werner Albert; Martin, Steven Michael

SOURCE: PCT Int. Appl., 37 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

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APPLICATION NO.
    PATENT NO.
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                                          WO 1997-GB2907
                           19980430
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    AU 9747154
                      A1
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                      B2 ...
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                           20001018
                                        GB 1996-21859
                                                            19961021
PRIORITY APPLN. INFO.:
                                                        A
                                        WO 1997-GB2907
                                                            19971021
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GT

A 5,6-dihydro- α -pyrone I (R = CO2H or CH3 and R1, R2 = H; or R = AB CO2H and one of R1 and R2 is H and the other is OH; or when R is CO2H, a pharmaceutically or veterinarily acceptable salt thereof) are provided. Processes for producing these compds., and their use as cytokine production inhibitors, are also described. Fermentative production and preparative esterification are included.

IC ICM C07D309-32

C12P007-40; C12P017-06; C07C059-42; C07C057-03; C12N001-14; C12N001-14; C12R001-645; C12R001-79

CC1-7 (Pharmacology)

Section cross-reference(s): 16, 27, 63

28921-94-0P 138875-82-8P 207220-97-1P TT

> RL: PUR (Purification or recovery); RCT (Reactant); PREP (Preparation); RACT (Reactant or reagent)

(reaction; 5,6-dihydro- α -pyrone cytokine production inhibitors,

production and preparation, and therapeutic use)

IT 28921-94-0P 138875-82-8P 207220-97-1P

RL: PUR (Purification or recovery); RCT (Reactant); PREP (Preparation); RACT (Reactant or reagent)

(reaction; 5,6-dihydro-α-pyrone cytokine production inhibitors, production and preparation, and therapeutic use)

RN28921-94-0 CAPLUS

2H-Pyran-2-one, 5,6-dihydro-5-hydroxy-6-(1E)-1-propenyl-, (5S,6S)- (9CI) CN(CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

RN 138875-82-8 CAPLUS

CN 2,4-Dodecadienoic acid, 4,6-dimethyl-, (2E,4E,6S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry as shown.

RN 207220-97-1 CAPLUS

CN 2-Propenoic acid, 3-[(2S,3S)-3,6-dihydro-3-hydroxy-6-oxo-2H-pyran-2-yl]-, (2E)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT